

# Integration of responses within and across *Arabidopsis* natural accessions uncovers loci controlling root systems architecture

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Phenotypic plasticity is presumed to be involved in adaptive change toward species diversification. We thus examined how candidate genes underlying natural variation across populations might also mediate plasticity within an individual. Our implementation of an integrative “plasticity space” approach revealed that the root plasticity of a single *Arabidopsis* accession exposed to distinct environments broadly recapitulates the natural variation “space.” Genome-wide association mapping identified the known gene *PHOSPHATE 1* (*PHO1*) and other genes such as *Root System Architecture 1* (*RSA1*) associated with differences in root allometry, a highly plastic trait capturing the distribution of lateral roots along the primary axis. The response of mutants in the Columbia-0 background suggests their involvement in signaling key modulators of root development including auxin, abscisic acid, and nitrate. Moreover, genotype-by-environment interactions for the *PHO1* and *RSA1* genes in Columbia-0 phenocopy the root allometry of other natural variants. This finding supports a role for plasticity responses in phenotypic evolution in natural environments.

GWAS | morphometrics | GxE interaction | QTL | RootScape

A long-standing debate in evolutionary biology is the relevance of phenotypic plasticity as a mechanism leading to species diversity (1). It has been argued that selection on plasticity responses to environment pressures could underlie fixed phenotypic changes between natural variants (2), providing a potentially rapid mechanism of evolutionary change (3). Thus, we tested the hypothesis that genes that enable a functional response to the environment within a population also underlie adaptive changes across natural variants. *Arabidopsis thaliana* offers ample opportunity to study genes involved in phenotypic plasticity in response to experimental laboratory perturbations, whereas its natural variants offer the opportunity to study the genetic basis for developmental variation observed in nature. We took a unique approach to integrating results from these two perspectives to uncover the molecular basis underlying individual plasticity and variation among natural variants. We focused on the root architectural system because it shows a high degree of plasticity under diverse environmental conditions (4–9). We used a quantitative phenotyping model to capture and integrate plastic changes in root system architecture in response to a range of experimental treatments within the laboratory reference accession Columbia-0 (Col-0). We next cross-referenced this plasticity space derived for an individual accession (Col-0) to the range of phenotypic differences in root architecture observed across *Arabidopsis* accessions that represent the extent of natural variation under one condition. This allowed us to identify candidate genes underlying root systems architecture using genome-

wide association mapping and to test their role in individual plasticity using mutants.

## Results

**Integrative Characterization of Phenotypic Plasticity and Natural Variation in Root Architecture.** To characterize the root plasticity “space” within an accession, we grew the *A. thaliana* laboratory strain Col-0 under a set of treatments using nutrients and hormones known to mediate different aspects of root development. This uncovered a range of plasticity within a single genotype across multiple treatments: control (KCl), auxin [Indol acetic acid (IAA)], cytokinin (CK), abscisic acid (ABA), nitrate (KNO<sub>3</sub>), and ammonium chloride (NH<sub>4</sub>Cl) (10) (Fig. 1A). Next, we characterized natural variation of root system architecture in 69 genotyped *Arabidopsis* accessions (12) grown in a single environment (KNO<sub>3</sub>) under which they exhibit a large breadth of natural phenotypic root variation (Fig. 1B). Root systems architecture was quantified holistically with a method (*RootScape*) that uses landmark-based morphometrics and has the advantage of being blind to the relevance of conventional morphological characters (11, 13). This method uses a 20-point landmark template that consists of a set of reference points fixed to developmental landmarks such as the base of the primary root and to points that capture plasticity to a greater degree, such as the end of the

## Significance

Species display a range of plastic phenotypes that presumably have evolved as a result of adaptation to heterogeneous environments. We asked whether the genetic mechanisms that underlie adaptation across populations also determine the response of an individual plant to environmental cues in *Arabidopsis*. Using an integrative root phenotyping approach, genes that underlie natural variation in root architecture across populations were shown to control plasticity responses within an individual. Together, our results uncover a genetic mechanism underlying the phenotypic plasticity of an individual and phenotypic diversity across natural variants.

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The authors declare no conflict of interest.

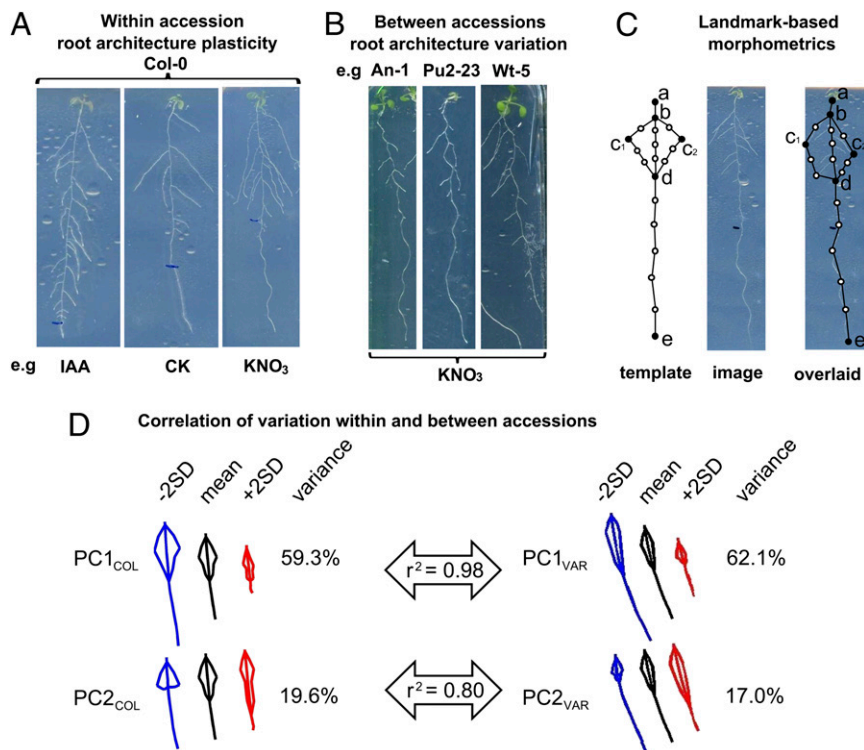
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**Fig. 1.** Root architecture plasticity within an accession (Col-0) broadly recapitulates natural variation quantified across 69 accessions. (A) Root phenotypes of Col-0 plants were grown under five conditions: IAA, CK, ABA, KNO<sub>3</sub>, and NH<sub>4</sub>Cl plus a control (KCl); roots of three treatment conditions are shown. (B) Root variation between 69 accessions grown under one condition (KNO<sub>3</sub>). (C) Landmark template to capture the root system architecture (11). Primary landmarks (black circles) are defined according to corresponding features in all roots; secondary landmarks (white circles) are evenly spaced between primary landmarks (11). (D) Two PCs capture more than 75% of the variation both within Col-0 (PC<sub>COL</sub>) and between accessions (PC<sub>VAR</sub>). PC1<sub>COL</sub> and PC2<sub>COL</sub> have high correlation to PC1<sub>VAR</sub> and PC2<sub>VAR</sub>, respectively. PC1<sub>COL</sub> is mainly a size effect, whereas PC2<sub>COL</sub> captures mainly root allometry, the length and distribution of lateral roots along the primary root. SD, standard deviation.

primary root, the widest lateral roots, and the end of the lateral roots. Intermediate points are added evenly between the fixed landmarks. In this way the template captures the main features of root architecture as an integrated system (11) (Fig. 1C).

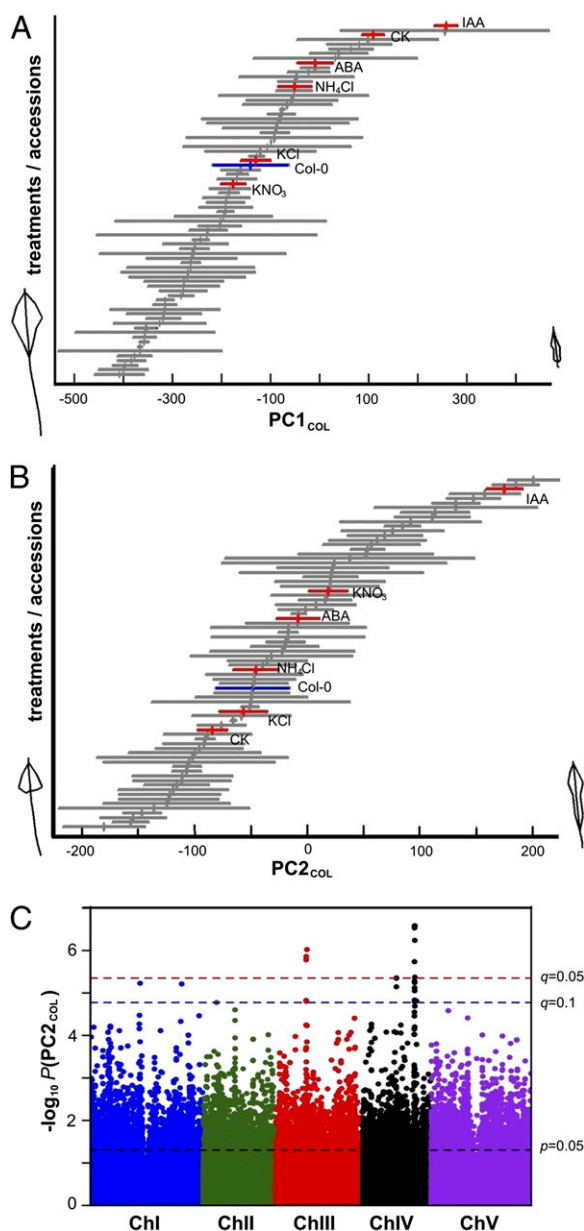
To create a framework to compare natural variation and plasticity responses, we used data from the landmark-based morphological root models and ran Principal Component Analyses (PCA). The resulting PCs uncovered the main trends of phenotypic plasticity in roots of the laboratory reference accession Col-0 (PC<sub>COL</sub>) and among the *A. thaliana* natural VARIANTS (PC<sub>VAR</sub>). In both root models (PC<sub>COL</sub> and PC<sub>VAR</sub>), PC1 and PC2 covered more than 75% of the variation in root systems architecture (Fig. 1D). PC1<sub>COL</sub> captures a size effect; PC2<sub>COL</sub> is an orthogonal axis to PC1<sub>COL</sub> and therefore captures a type of variation unrelated to size. Thus, PC2<sub>COL</sub> seems to capture the proportion of the primary root that contains visible lateral roots, in other words, the distribution of lateral roots in the proximal–distal axis regardless of root size (Fig. 1D). Similarly, a PCA of the VAR space across 69 natural *Arabidopsis* accessions captures a size-effect trend of the variation (PC1<sub>VAR</sub>) and the lateral root allometry effect (PC2<sub>VAR</sub>). Overall, the PC analyses suggest that the variation captured within an individual accession (PC<sub>COL</sub>) is similar to the range of variation observed in natural accessions (PC<sub>VAR</sub>) under our growth conditions (Fig. 1D). This suggests a shared genetic mechanism controlling both phenotypic plasticity and natural variation, which would imply an important role of plasticity in adaptive change.

The similarity of variation in root systems architecture observed between the plasticity space of Col-0 and across the natural accessions—(PC1<sub>COL</sub> ~ PC1<sub>VAR</sub>) and (PC2<sub>COL</sub> ~ PC2<sub>VAR</sub>)—allowed us to use these PC axes as phenotypic metrics to perform quantitative comparisons. We projected the PC values for both datasets COL and VAR into the PC1<sub>COL</sub> and PC2<sub>COL</sub> axes (reference plasticity space) (Fig. 2). Surprisingly, the laboratory reference accession Col-0 (Fig. 2A and B, blue bar), which has been historically selected for other purposes, localized roughly to the middle of the phenotypic axes of both PCs, corresponding in root phenotype to a “mean accession” relative to the other natural

variants (Fig. 2A and B, gray and blue bars). However, the treatments (Fig. 2A and B, red bars) expanded the phenotypic breadth of Col-0, pushing its plasticity mainly toward the higher end of the PC<sub>COL</sub> axes, relative to the mean. This distribution suggests underlying developmental constraints for the plasticity to explore the lower end of the PC<sub>COL</sub> space. We tested this using other *Arabidopsis* accessions and found that their plasticity is mostly directed toward the higher end of the PC axis (i.e., PC2<sub>COL</sub>) (Fig. S1). PC2<sub>COL</sub> captures the distribution of the lateral roots on the primary roots. The lower end shows a narrower distribution on a range of accessions (i.e., Tamm-27) and Col-0 plasticity phenotypes (i.e., CK). The absence of phenotypes with lower PC2<sub>COL</sub> might reflect a bias of our selected treatments.

The projection onto the PC1<sub>COL</sub> and PC2<sub>COL</sub> phenotypic spaces (Fig. 2A and B) also allowed us to establish that the distribution of Col-0 phenotypes resulting from experimental treatments (Fig. 2A and B, red bars) covers ~50% of the distribution of phenotypes exhibited by the natural accessions along PC1<sub>COL</sub> and ~65% of the distribution in PC2<sub>COL</sub> (Fig. 2A and B, gray and blue bars). We performed the reciprocal analysis and projected the same datasets into the PC1<sub>VAR</sub> and PC2<sub>VAR</sub> axes (“natural variation space”) obtaining similar results (Fig. S2). A pair-wise comparison of the phenotype projections showed that PC1<sub>COL</sub> and PC2<sub>COL</sub> are highly correlated to PC1<sub>VAR</sub> and PC2<sub>VAR</sub>, respectively (Fig. 1D). Thus, the plasticity space of individual *Arabidopsis* accessions recapitulates natural variation in root architecture when exposed to distinct laboratory-induced physiological environments.

**Identification of Candidate Genes Underlying Natural Variation in Root Architecture.** The projections of the PC values from the natural variants into the PC1<sub>COL</sub> and PC2<sub>COL</sub> phenotypic space were next used to map genes underlying both natural variation and phenotypic plasticity. To do this, we looked for associations between the PC1<sub>COL</sub> or PC2<sub>COL</sub> phenotypes exhibited in the natural accessions and the 214,000 SNP dataset of polymorphisms in *Arabidopsis* (14). No significant genome-wide association study (GWAS) associations were identified for the PC1<sub>COL</sub> trait (overall root size), suggesting that size might be



**Fig. 2.** Root plasticity variation within an accession (Col-0) spans a range of plasticity exhibited by *Arabidopsis* natural variants and maps to regions in chromosome III and IV. (A and B) Bars indicate SE. Red bars, Col-0 in the five treatments IAA, CK, ABA, KNO<sub>3</sub>, and NH<sub>4</sub>Cl and a KCl control ( $n = 20$ ); gray bars, phenotypes of 69 *Arabidopsis* accessions grown under a single (KNO<sub>3</sub>) condition ( $n = 3$ –4); blue bars, reference Col-0 accession. Morphometrics modeled root systems architecture phenotypes; those corresponding to extreme root PC1<sub>COL</sub> and PC2<sub>COL</sub> phenotypes are shown. (C) Manhattan plot illustrating the GWAS mapping of the PC2<sub>COL</sub> phenotype.

controlled by many loci of small effect. However, the PC2<sub>COL</sub> trait (allometry of lateral roots) revealed 12 significant GWAS associations at the false-discovery rate (FDR) level of  $q < 0.05$  or 19 associations at  $q < 0.1$ . Most of these SNPs mapped to two genomic regions in chromosomes III and IV (Fig. 2C), partially overlapping regions with previously identified root morphology quantitative trait loci (QTL) (4–6, 15–18). We assumed a 5-kb window before the first and after the last significant SNPs, given that linkage disequilibrium decays more than 50% in *Arabidopsis* in that range (19). Thus, the region in chromosome (Chr) III had five SNP associations in a 212-kb interval (Chr III:

8181817–8393525 bp) containing 49 genes (Table S1, Chromosome III region). The region in chromosome IV had 18 significant SNPs in a 112-kb interval (Chr IV: 13988547–14100946 bp) containing 39 genes (Table S1, Chromosome IV region). We focused on a subset of 11 genes that each contain highly significant SNP associations (FDR:  $q < 0.05$ ) (Table S2). Among these candidate genes is *PHOSPHATE 1* or *PHO1* (At3g23430) shown previously to play a role in inorganic phosphate loading into the xylem and aspects of root development (20, 21). Several other uncharacterized genes were also identified.

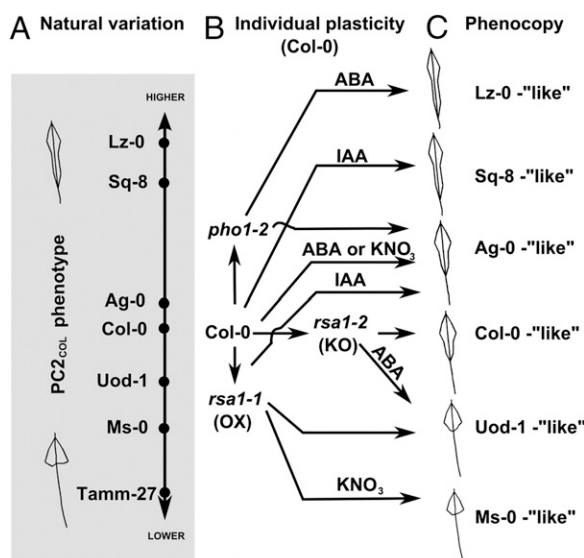
To test whether the corresponding mutant alleles show phenotypic effects for PC2<sub>COL</sub>, we characterized the root phenotypes in mutants of six candidate genes in the chromosome III and IV regions (Fig. 3A). *pho1-2* (22) showed a significantly lower PC2<sub>COL</sub> phenotype than wild-type Col-0 (Fig. 3A), supporting the hypothesis that *PHO1* is one of the underlying QTL in chromosome III. A second gene that showed a significant phenotype in PC2<sub>COL</sub> is At4g28410 or *Root System Architecture 1* (*RS1*). *RS1*, a gene identified in this screen, encodes a protein that belongs to the tyrosine transaminase family, with similarity to *SUPERROOT 1* (*SUR1*; At2g20610) (23), a gene involved in glucosinolate biosynthesis from tryptophan derivatives (24).

Among other defects, the *sur1* mutant overproduces auxin because the conversion to glucosinolates is blocked, which canalizes tryptophan derivatives toward auxin production, thus exhibiting extensive proliferation of lateral roots (25). We thus postulate that the *SUR1*-like gene, *RS1*, has a redundant function to *SUR1*, and loss-of-function mutations (*rs1-2* = At4g28410-KO) are unlikely to show strong phenotypic effects; however, overexpression alleles (*rs1-1* = Atg428410-OX) would be expected to have an auxin deficiency-like phenotype, consistent with the phenotype of the *rs1-1* gain-of-function mutant (Fig. 3A and B). *rs1-1* is a tDNA insertion mutant mapping to the 5' UTR of At4g28410, whereas *rs1-2* is a tDNA insertion mapping to 969 bp from the start codon, producing a predicted truncated protein of 327 amino acids (full length is 447 amino acids). To test the possible role of *RS1* in auxin homeostasis, we obtained the genome-wide expression profile of the *rs1-1* overexpression mutant. We found that 104 ( $P < 0.001$ ) genes or 492 ( $P < 0.01$ ) genes are more than 1.5-fold differentially expressed compared with the sibling wild type (Table S3). We compared the list of misregulated genes in the *rs1-1* mutant to a list of 3,186 genes regulated by the synthetic auxin naphthalene-1-acetic acid in the root (26). We found significant over-representations of the intersections of auxin-regulated genes with genes misregulated in the *rs1-1* mutant for both of our datasets ( $P < 0.001$ ), compared with random generated gene lists of the same size (27, 28), further suggesting a *rs1-1* role in auxin homeostasis. To further test whether expression of *RS1* mRNA is causative to changes in root allometry, we tested whether natural variation in *RS1* activity is correlated with changes in PC2<sub>COL</sub>. To do this, we obtained normalized *RS1* expression in 10 *Arabidopsis* accessions (Fig. S3) and compared them to their respective PC2<sub>COL</sub> phenotypes. Consistent with the result on the *rs1-1* overexpression allele, we found a negative correlation between *RS1* mRNA levels and the PC2<sub>COL</sub> phenotype (Pearson correlation:  $-0.55$ ). This correlation of gene activity and phenotypic changes in PC2<sub>COL</sub> in the *rs1-1* mutant, and the correlation between *RS1* expression and PC2<sub>COL</sub> phenotypes in the natural accessions also supports the notion that *RS1* (At4g28410) underlies the QTL for PC2<sub>COL</sub> that maps to chromosome IV.

Using the *Arabidopsis* 1,001 Genomes browser (1001genomes.org), we identified SNP positions in the *RS1* and *PHO1* loci. For *RS1*, the SNP at position 14051900 in chromosome IV is 348 bp upstream of the 5' UTR. This opens the possibility that *cis*-regulatory variation at *RS1* is driving variation in root allometry that is supported by our studies of *RS1* expression in the *rs1-1* mutant (Fig. 3A and B) and in natural variants (Fig. S3). For *PHO1*, we found that the SNP at position 8388425 in chromosome III (TAIR 8 annotation) is within an exon of the coding region. Furthermore, the SNP is nonsynonymous; a cytosine







**Fig. 4.** The *PHO1*-*RSA1*-dependent  $G \times E$  interaction expands the root plasticity space of Col-0 to phenocopy a range of natural accessions. (A) Axis representing natural variation on  $PC2_{COL}$  phenotype with examples of accession's positions and root allometry phenotypes. (B) Treatments with  $KNO_3$ , ABA, or IAA in *rsa1-1* gain-of-function (OX) and *rsa1-2* loss-of-function (KO) mutants have contrasting effects to Col-0. (C) Col-0 and *rsa1* mutant alleles treated with  $KNO_3$ , ABA, or IAA phenocopy the root architecture of natural variants.

allometry in the Col-0 accession in turn phenocopies the root allometry trait of a range of natural accessions (Fig. 4C). This suggests that the mechanisms controlling phenotypic plasticity of root allometry Col-0 are similar to the ones controlling natural variation in *Arabidopsis* accessions. Moreover, the accessibility of the phenotypic plasticity space of an individual accession depends on the activity of genes controlling natural variation, as represented by *RSA1* and *PHO1*. These findings reopen the discussion of the relevance of phenotypic plasticity genes in natural variation and adaptive evolution (1–3, 29).

Our study also highlights a strategy for intersecting experimentally perturbed phenotypes within a laboratory strain with those observed across a panel of natural variants to uncover genetic mechanisms underlying plasticity and natural variation for any evolved trait of interest. Contextualizing root plasticity within laboratory variants and natural variation in accessions under laboratory conditions is a powerful strategy to dissect the influence of factors on plasticity typically composed of complex multigenic traits, such as the impact of hormones on root system architecture. In developing a method to cross-reference the genes involved in phenotypic plasticity and natural variation, we open a window to understanding the possible mechanisms that characterize adaptive change in *Arabidopsis*.

## Methods

**Growth Conditions and Phenotyping.** Seeds were disinfected with a solution of ethanol, bleach, and water 4:1:3 and three rinses of sterilized water. Seeds were sown on square plates of custom-made MS media (–sucrose/–nitrogen) supplemented with sucrose 0.1%, nitrogen (as indicated below), MES sodium salts 0.05% (Gibco BRL), and agar 1% (Bacto Agar BD). Plates were kept at 4 °C for 4 d in the dark and then placed vertically in a growth chamber under 22 °C, long-day 16h/8h condition at  $125 \mu\text{mol}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$  light intensity (Percival Scientific). For the phenotypic characterization of accessions and mutants, the custom MS media was supplemented with 5 mM of  $KNO_3$  and 1% of sucrose as carbon source, grown at  $50 \mu\text{mol}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$  light intensity; roots were imaged 16 d after sowing. For the IAA, CK, ABA,  $KNO_3$ ,  $NH_4Cl$ , and control (KCl) treatments in Col-0 and mutants, the seeds were sown on media containing 1 mM of KCl, and seedlings were transferred to the fresh plates containing 500 nM of IAA, 500 nM of kinetin (CK), 1  $\mu\text{M}$  of ABA, 1 mM of  $KNO_3$ , 1 mM of  $NH_4Cl$ , or 1 mM KCl (control), and roots were imaged 14 d after germination (4 d after transfer to treatment plates).

Plate images were obtained at 300-dpi resolution using a scanner. Landmark data and morphometric analysis were obtained using the software "Shape Model Toolbox" (12) in Matlab as described (13). Before analyzing the root variation, procrustes for translation and rotation were applied to the datasets (11, 13). This was done by aligning the datasets according to their centroid and then rotating them about the centroid to minimize the distance between corresponding landmarks; no scaling was applied. PCA was carried out on the covariance matrix to obtain the main trends of the variation (PCs), which were used as traits. Projections of other datasets were used to quantify phenotypes according to the corresponding  $PC_{COL}$ . For details on the PCA methods and projections, see refs. 13 and 35. Landmark datasets to generate the models are provided in Dataset S1.

**Sampling and Statistical Tests.** The phenotypic plasticity space  $PC_{COL}$  was created by growing Col-0 in the treatments described above with 20 plants per treatment. The accession data were obtained by growing three to four plants of each of the 96 Nordborg lines. Only 69 were phenotyped with the RootScape method (11), as the rest did not outgrow visible lateral roots under our conditions. To characterize the mutant phenotypes of candidate genes, homozygote tDNA insertion mutants and wild-type siblings were identified using primers obtained using default settings on the iSect tool (<http://signal.salk.edu/tdnaprimers.2.html>). Plates were sown side by side with the mutant and wild-type siblings to account for background and plate effects. The dataset consisted of  $n = 239$  plants of Col-0; 36 of *pho1-2*; 23 of GABI\_297C02; 25 of GABI\_764B10; 24 of SAIL\_47\_B10; 27 of SALK\_094117; 22 of GABI\_288A12; 31 of WiscDsLoxHs186\_11; 22 of SALK\_011695C; and 22 of SALK\_094441C. The projected  $PC2_{COL}$  data were analyzed according to the following REML mixed model:  $PC2_{COL} = \beta_{\text{genotype}} + \beta_{\text{plate}} + \varepsilon$  (Fig. 3A). To characterize the plasticity response of the *RSA1*/At4g28410 mutants, a minimum of 11 plants per treatment per genotype were analyzed, for a total of 291 plants. This dataset was analyzed together with the data to generate the  $PC_{COL}$  space (above).  $G \times E$  interactions were tested according to the following two-way ANOVA model:  $PC2_{COL} = \beta_{\text{genotype}} + \beta_{\text{treatment}} + \beta_{\text{genotype} \times \text{treatment}} + \varepsilon$  (Fig. 3 C–E). To further test *pho1-2* plasticity in a phosphate-depleted environment, we custom-made MS media following the manufacturer's recipe, but also substituted the moles of  $PO_4$  in  $KH_2PO_4$  for KOH and imaged the roots 4 d after transfer to phosphate-depleted media.

**Genome-Wide Association Study.** After standardizing each  $PC_{COL}$  root trait to a mean of zero and a SD of 1, we performed genome-wide association mapping using the SNP database from Atwell et al. (14), which documents over 214,000 SNPs (an average of 1 SNP/500 bp) in 69 different inbred lines from the wild. We filtered the database to include only SNPs with a minor allele frequency greater than 0.10, which left 177,623 SNPs for mapping.

To account for genome-wide patterns of relatedness that can confound the results of GWAS studies (36), we used all 214,000 SNPs to construct a similarity matrix representing the proportion of loci that is identical in state between any pair of lines ( $K$ ; 36). For each trait, we then separately fit the model  $y = X\alpha + Zu + \varepsilon$ , where  $y$  is a vector of phenotypes,  $X$  is a matrix of single-locus genotypes,  $\alpha$  is a vector of allele effects to be estimated,  $Z$  is an identity matrix,  $u$  is a matrix of random deviates due to genome-wide relatedness (as inferred from  $K$ ), and  $\varepsilon$  is a vector of residual errors. The analysis was conducted using the EMMA approach (37) in R version 3.0.1 (R Development Core Team 2013). To account for multiple simultaneous tests (because  $\alpha$  is modeled separately for each SNP), we calculated  $P$  values that were adjusted for the genome-wide FDR ( $q$ -values) using the  $q$ -value package (38) in R (R Core Team 2013) (39).

**Gene Expression Analysis.** For expression analysis using quantitative PCR, the RNA extraction were carried out on roots collected from a *rsa1-2* mutant and sibling wild type at day 12, grown side by side on vertical agar plates, 5 mM of  $KNO_3$ , and 1% of sucrose at  $50 \mu\text{mol}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$  light intensity. For each of the three replicates, we pooled tissue from three roots. For assays in accessions, plants were grown in liquid media in phytotrays, and total RNA was extracted using RNeasy minikit (Qiagen). Double-stranded cDNA was synthesized by the SuperScript RT-PCR system (Invitrogen). PCR were performed using the LightCyclerFastStart DNA master<sup>PLUS</sup> SYBR Green I (Roche) in a LightCycler 480 (Roche). Expression level of At4g28410 was quantified using the oligos 5'-GTGGTGATAATGAATCCTCACAAC-3' and 5'-CCATCGGG-ACAAATTTATTCTCT-3'. Five standard reference genes were used to quantify relative expression: *Clathrin*/At4g24550 (5'-AGCATACACTGCGTGCAAAG-3' and 5'-TCGGCTGTGCACATATCTC-3'), *ACT2*/At3g18780, At1g49240 (5'-GGTACATGTGTCTCAGGGTGG-3' and 5'-AACGACCTTAATCTTCATGCTGC-3'), *SIP1A*/At3g04090 (5'-TCCTTGTCATTGTTAGATCCACAC-3' and 5'-TAAATGTTCTCAACCGGAAGAGAGTC-3'), *ATJ3*/At3g44110 (5'-TCCAACCAATTTGTCTCTTGCT-3' and 5'-TAAATGTTCTCAACCGGAAGAGAGTC-3').

and 5'-AACAAAGTTTCGATGTTCCACC-3'), and *UBI10*/At4g0532 (5'-GGCCTTG-TATAATCCCTGATGAATAAG-3' and 5'-AAAGAGATAACAGGAACGGAACA-TAGT-3'). All PCRs were performed with annealing of 60 °C. PCR efficiency of At4g28410 was tested with a standard curve in each plate, using four serial dilutions of a wild-type sample: 1/1, 1/10, 1/100, and 1/1000. For genome-wide expression in the *rsa1-1* mutant allele ( $n = 3$ ) and the sibling wild type ( $n = 3$ ), we used standard protocols from Affymetrics to amplify, label, and hybridize RNA samples to the ATH1 Affymetrics chip. Raw data were processed in MASv5.0 and two-tailed  $t$  test was performed in R (R Core Team 2013) (40).

**Landscape Genetic Analysis.** We took the significant SNP in *PHO1* (At3g23430) and the significant SNP near *RSA1* (At4g28410) and examined how they were structured on the landscape as a function of several environmental factors. We used a list of accessions and geographic coordinates of their origins (41). We intersected this list with lists of genes for which SNP genotype information is available at the two SNP loci of interest (14). We limited this list to accessions falling within the native range of *A. thaliana* in Europe and Asia (−11° to 86° E and 35° to 71° N), as estimated based on ref. 42. We then filtered this list by the “red list” and the “yellow list” of putative misidentified accessions previously flagged (41). A total of 726 accessions remained after intersecting and filtering (Fig. S5A and Table S5).

Separately for each SNP, we examined how the SNP was structured on the landscape as a function of temperature and humidity variables used in previous studies (Table S4). All environmental factors were at a 10 arc-minute (~13 km) scale worldwide. We extracted the values of the environmental factors at the locations origin of the accessions using ArcMap version

9.3 (Esri Inc.). We then thinned the list of environmental factors to those with correlation coefficients less than |0.6| among the locations of the accessions (Table S4).

The logistic regression model was based on the logistic function  $y = 1/(1 + e^{-z})$ , where  $y$  is a binary dependent variable (one SNP allele or the other); and  $z = \beta_0 + \beta_1 S_1 + \dots + \beta_x S_x + \beta_1 E_1 + \dots + \beta_x E_x + \varepsilon$ , where  $\beta_0$  is the intercept,  $S_1 - S_x$  are the covariates to account for spatial autocorrelation (Table S4),  $E_1 - E_x$  are the environmental variables,  $\varepsilon$  is the residual error, and the  $\beta$  terms are the corresponding regression coefficients. The analysis was performed using the lrm function of the rms package (39) in R (R Development Core Team 2013) (40).

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